# CLINICAL LABORATORY IMPROVEMENT AMENDMENTS OF 1988 (CLIA) APPLICATION FOR CERTIFICATION

I. GENERA	LINFORMATION			
<ul><li>☐ Initial Application</li><li>☐ Change in Certification Type</li></ul>	CLIA Identification Number     (If an initial application leave blank, a number will be assigned)			
Facility Name	Federal Tax Identification Number			
	Telephone No. (Include area code) Fax No. (Include area code)			
Facility Address — Physical Location of Laboratory (Building, Floor, Suite if applicable.)	Mailing/Billing Address (If different from street address, include attention line and/or Building, Floor, Suite)			
Number, Street (No P.O. Boxes)	Number, Street			
City State Zip Code	City State Zip Code			
Name of Director				
Last First Middle initial				
II. TYPE OF CERTIFIC	ATE REQUESTED (Check One)			
<ul> <li>□ Certificate of Waiver (Complete Sections I – VI</li> <li>□ Certificate for Provider Performed Microscopy</li> <li>□ Certificate of Complete Sections I</li> </ul>	Procedures (PPMP) (Complete Sections $I - X$ )			
☐ Certificate of Accreditation (Complete Sections organization(s) your laboratory is accredited by applied for accreditation for CLIA purposes	I through X) and indicate which of the following y for CLIA purposes, or for which you have			
$\square$ JCAHO $\square$ AOA	☐ AABB			
□ CAP □ COLA	ASHI			

According to the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number. The valid OMB control number for this information collection is 0938-0581. The time required to complete this information collection is estimated to average 30 minutes to 2 hours per response, including the time to review instructions, search existing data resources, gather the data needed, and complete and review the information collection. If you have any comments concerning the accuracy of the time estimate(s) or suggestions for improving this form, please write to: CMS, 7500 Security Boulevard, N2-14-26, Baltimore, Maryland 21244-1850.

III. TYPE OF LABORATORY (Check the one most descriptive of facility type)								
01 Ambulator	01 Ambulatory Surgery Center 09 Hospice				17	17 School/Student Health Service		
02 Community	_ 02 Community Clinic 10 Hospital					18 Skilled Nursing Facility/Nursing Facility		
03 Comp. Outpatient Rehab. Facility 11 Independent						19 Physician Office		
	esting Site in Health	-	12 Industrial		20	20 Other Practitioner (Specify)		
-	Renal Disease Dialy		13 Insurance			21 Tissue Bank/Repositories		
06 Health Fair	-			re Fac. for Mentally R		Blood Banks		
	in. Organization		15 Mobile Laborato	· · · · · · · · · · · · · · · · · · ·		Rural Health Clinic		
	•			ıy			Lloolth Contor	
08 Home Hea	iith Agency		16 Pharmacy			24 Federally Qualified Health Center		
	icare/Medicaid cert	, —			25 Ambulance			
If yes, indica	te Medicare provide				26	26 Public Health Laboratories		
	Medica	id number			27	Other		
							•	
1	V. HOURS OF	LABORATORY	Y TESTING (Li	st times during w	hich <b>laboratory</b>	testing is perfor	rmed)	
	SUNDAY	MONDAY	TUESDAY	WEDNESDAY	THURSDAY	FRIDAY	SATURDAY	
FROM: AM								
H	<b></b>							
TO: AM	<u> </u>			<u> </u>	<u> </u>	J		
PM								
/E 1:1	1	1				1		
(For multiple sit	tes, attach the add	ditional informat	ion using the san	ne format.)				
	V. MULTIPL	E SITES (must	meet one of the	regulatory excep	otions to apply f	or this provision	1)	
Are vou annis	ying for the mul	tinle site evcenti	ion?					
□ No If <b>no</b> ,	go to section VI.		• •	al number of site	s under this cert	ificate	and complete	
		re	emainder of this	section.				
	Indicate wh	ich of the follow	ving regulatory	exceptions appli	es to vour facili	ty's operation		
In this a ma					•	-		
	t-for-profit or Fed ngaged in limited (1						ated at contiguous	
	mplexity or waived				buildings on the same campus within the same physical location or street address and under common direction that is filing for a			
	iling for a single c				single certificate for these locations? $\square$ Yes $\square$ No			
☐ Yes ☐	No			_	If yes, list name or department, location within hospital and			
If yes, list na	me, address and te	sts performed for e	ach site below.		specialty/subspecialty areas performed at each site below.			
-		-				•		
If add	ditional space is	needed, check l	here and a	ttach the additio	onal information	n using the sam	e format.	
NAME AND ADDRESS / LOCATION			TEST	rs performed /	SPECIALTY / SU	BSPECIALTY		
Name of labor	atory or hospita	al department						
	,	•						
Address/location (number, street, location if applicable)								
Address/location (number, street, location if applicable)								
0'' 0' ' 715			T.I I NI.					
City, State, ZIF	J		Telephone No.					
Name of laboratory or hospital department								
Address/location (number, street, location if applicable)								
City, State, ZIP Telephone No.								
( )								
Name of laboratory or hospital department								
Address/location	n (number, stree	t location if appli	icable)					
/ wuress/iocallo	ii (Hairiber, Sude	., юсанын п аррп	odbio <sub>j</sub>					
0:1 0: : =:-		Т	T.1 **					
City, State, ZIF	,		Telephone No.					
			( )					

Indicate the estimated TOTAL	ANNUAL TES	T volume for all	waived	l tests performed.		
	VII. NON	VAIVED TESTI	NG (I	icluding PPMP testing)		
If you perform testing other the nultiple sites, the total volume				te the information below.	If applying for (	one certificate for
Place a check $()$ in the box present volume for each specialty. <b>D</b> quality assurance or proficient information included with the appropriate of a subspecialty for which you are	o not include test cy testing when oplication packag ccreditation, ind	ting not subject to calculating test e.)	CLIA volum	a, waived tests, or tests run e. (For additional guidance accreditation organization	for quality cone on counting test	trol, calculations, st volume, see the blicable specialty/
SPECIALTY / SUBSPECIALTY	ACCREDITING ORGANIZATION	ANNUAL TEST VOLUME			ACCREDITING ORGANIZATION	ANNUAL TEST VOLUME
Histocompatibility	ORGANIZATION			Hematology		
Transplant				Immunohematology		
Nontransplant				ABO Group & Rh Group		
Microbiology				Antibody Detection		
Bacteriology  Mycobacteriology				(transfusion)		
<ul><li>Mycobacteriology</li><li>Mycology</li></ul>				Antibody Detection		
Parasitology				(nontransfusion)		
☐ Virology				Antibody Identification		
				Compatibility Testing		
Diagnostic Immunology  Syphilis Serology						
				Pathology Histopathology		
General Immunology				Oral Pathology		
Chemistry				Cytology		
Routine				Cytology		
Urinalysis				Radiobioassay		
Endocrinology						
Toxicology				Clinical Cytogenetics		
		l				
TOTAL ES	<b>FIMATED A</b>	NNUAL TEST	r vo	LUME		_

VI. WAIVED TESTING

	VIII. TYP	E OF CONTROL		
Enter the appropris			(Futor only one code)	
	-	the list below	(Enter only one code)	
•	For Profit	Government	08 Federal	
<u>-</u>	04 Proprietary	05 City 06 County	09 Other Government	
02 Private		07 State	(Spec	cify)
03 Other(Specify)		07 State	V-T	37
IX. DIRI	ECTOR AFFILIATIO	N WITH OTHER LAB	ORATORIES	
If the director of this laboratory serves a following:	as director for additiona	l laboratories that are sepa	rately certified, please complete the	
NAME OF LABORATORY		ADDRESS	CLIA IDENTIFICATION N	UMBER
W TAIL		ED IN LABORATORY	THE COUNTY OF TH	
Indicate the total number of individuals				
individuals who only collect specimens				
<b>highest</b> laboratory position in which they		thologist serves as director	, technical supervisor and general sup	ervisor.
This individual would only be counted or	ince (under director).)			
A. WAIVED TESTING	•	D TESTING (including		
Total No. of Individuals	Total No. of	Individuals Director	Technical supervisor General supervisor	
	Clinica	consultant	Testing personnel	
		consultant	<b>8 F</b> • • • • • • • • • • • • • • • • • • •	
	Cytot	echnologist		
ATTENTION: READ THE	EOLL OWING C	ADEELII I V DEE	DE SIGNING ARRIVA	TION
Any person who intentionally violates a				
promulgated thereunder shall be imprison the conviction is for a second or subsequ				
or fined in accordance with title 18, Uni			nan be imprisoned for not more than	13 years
or rined in decordance with title 10, cm	ica states code of sour.			
Consent: The applicant hereby agrees th				
found necessary by the Secretary of Hea				
Act as amended. The applicant further a Secretary, to inspect the laboratory and it				
information or materials necessary to de				
compliance with CLIA requirements.	<i>,</i>	<i>y</i>	,	
			T	
SIGNATURE OF OWNER/DIRECTOR OF LA	BORATORY (Sign in ink)		DATE	

# THE CLINICAL LABORATORY IMPROVEMENT AMENDMENTS (CLIA) APPLICATION (FORM CMS-116)

#### INSTRUCTIONS FOR COMPLETION

CLIA requires every facility that tests human specimens for the purpose of providing information for the diagnosis, prevention or treatment of any disease or impairment of, or the assessment of the health of, a human being to meet certain Federal requirements. If your facility performs tests for these purposes, it is considered, under the law, to be a laboratory. CLIA applies even if only one or a few basic tests are performed, and even if you are not charging for testing. In addition, the CLIA legislation requires financing of all regulatory costs through fees assessed to affected facilities.

The CLIA application (Form CMS-116) collects information about your laboratory's operation which is necessary to determine the fees to be assessed, to establish baseline data and to fulfill the statutory requirements for CLIA. This information will also provide an overview of your facility's laboratory operation. All information submitted should be based on your facility's laboratory operation as of the date of form completion. **NOTE: WAIVED TESTS ARE NOT EXEMPT FROM CLIA. FACILITIES PERFORMING ONLY THOSE TESTS CATEGORIZED AS WAIVED MUST APPLY FOR A CLIA CERTIFICATE OF WAIVER.** 

THE GENERAL INFORMATION SECTION AND ALL APPLICABLE SECTIONS MUST BE COMPLETED. INCOMPLETE APPLICATIONS CANNOT BE PROCESSED AND WILL BE RETURNED TO THE FACILITY. PRINT LEGIBLY OR TYPE INFORMATION

# I. GENERAL INFORMATION

For an initial applicant, the CLIA identification number should be left blank. The number will be assigned when the application is processed.

Be specific when indicating the name of your facility, particularly when it is a component of a larger entity; e.g., respiratory therapy department in XYZ Hospital. For a physician's office, this may be the name of the physician. NOTE: The information provided is what will appear on your certificate.

Facility street address must be the <u>actual</u> physical location where testing is performed, including floor, suite and/ or room, if applicable. **DO NOT USE A POST OFFICE BOX NUMBER OR A MAIL DROP ADDRESS FOR THE NUMBER AND STREET OF THE ADDRESS.** If the laboratory has a separate mailing or billing address, please complete that section of the application.

#### II. TYPE OF CERTIFICATE REQUESTED

When completing this section, please remember that a facility holding a—

- Certificate of Waiver can only perform tests categorized as waived;\*
- Certificate for Provider Performed Microscopy Procedures (PPMP) can <u>only</u> perform tests categorized as PPMP, or tests categorized as PPMP and waived tests;\*
- Certificate of Compliance can perform tests categorized as waived, PPMP and moderate and/or high complexity tests provided the applicable CLIA quality standards are met; and
- Certificate of Accreditation can perform tests categorized as waived, PPMP and moderate and/or high complexity tests provided the laboratory is currently accredited by an approved accreditation organization.\*\*
- \*A current list of waived and PPMP tests may be obtained from your State agency. Specific test system categorizations can also be reviewed via the Internet on **www.cdc.gov/phppo/dls/.**
- \*\*If you are applying for a Certificate of Accreditation, you must include evidence of accreditation for your laboratory by an approved accreditation organization for CLIA purposes or evidence of application for such accreditation with the completed Form CMS-116.

#### III. TYPE OF LABORATORY

Select the type of laboratory designation that is most appropriate for your facility from the list provided. If you cannot find your designation within the list, contact your State agency for assistance.

## IV. HOURS OF ROUTINE OPERATION

Provide only the times when actual laboratory testing is performed in your facility.

# V. MULTIPLE SITES

You can only qualify for the multiple site provision (more than one site under one certificate) if you meet one of the CLIA regulatory exceptions outlined on the form.

# VI. WAIVED TESTING

Include only the estimated annual volume for those tests that are waived.

# VII. NON-WAIVED TESTING (Including PPMP)

Follow the specific instructions on page 3 of the Form CMS-116 when completing this section. (Note: The Accrediting Organization column should reflect accreditation information for CLIA purposes only; e.g., JCAHO, etc.).

#### VIII. TYPE OF CONTROL

Select the code which most appropriately describes your facility. Proprietary/for profit entities must choose "04".

#### IX. DIRECTOR AFFILIATION WITH OTHER LABORATORIES

List all other facilities for which the director is responsible. Note that for a Certificate of PPMP, Certificate of Compliance or Certificate of Accreditation, an individual can only serve as the director for no more than five certificates.

#### X. INDIVIDUALS INVOLVED IN LABORATORY TESTING

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\*

Once the completed Form CMS-116 has been returned to the applicable State agency and it is processed, a fee remittance coupon will be issued. The fee remittance coupon will indicate your CLIA identification number and the amount due for the certificate, and if applicable the compliance (survey) or validation fee. If you are applying for a Certificate of Compliance or Certificate of Accreditation, you will initially pay for and receive a Registration Certificate. A Registration Certificate permits a facility requesting a Certificate of Compliance to perform testing until an onsite inspection is conducted to determine program compliance; or for a facility applying for a Certificate of Accreditation, until verification of accreditation by an approved accreditation organization is received by CMS.

If you need additional information concerning CLIA, or if you have questions about completion of this form, please contact your State agency.

# TESTS COMMONLY PERFORMED AND THEIR CORRESPONDING LABORATORY SPECIALTIES/SUBSPECIALTIES

# **HISTOCOMPATABILITY**

HLA Typing (disease associated antigens)

# **SYPHILIS SEROLOGY**

**RPR** 

FTA, MHA-TP

#### **GENERAL IMMUNOLOGY**

Mononucleosis Assays Rheumatoid Arthritis Febrile Agglutins Cold Agglutinins

HIV

Antibody Assays (hepatitis, herpes, etc.)

**ANA** Assays

Mycoplasma pneumoniae Assays

# **PARASITOLOGY**

**Direct Preps** 

Ova and Parasite Preps

Wet Preps

#### **CHEMISTRY**

# **Routine Chemistry**

Albumin BUN
Ammonia Uric acid
Bilirubin, Total ALT/SGPT
Bilirubin, direct AST/SGOT

Calcium SGGT Chloride Alk Phos Cholesterol,total Amylase

CO2, total CPK/CPK isoenzymes

Creatinine CKMB

Glucose HDL Cholesterol

pH Iron pO2 LDH

pCO2 LDH isoenzymes
Phosphorous Magnesium
Potassium Ferritin
Protein,total Folic Acid
Sodium Vitamin B12

Triglycerides PSA

#### **Urinalysis**

Automated urinalysis

Urinalysis with microscopic analysis Urine specific gravity by refractometer Urine specific gravity by urinometer Urine protein by sulfasalicylic acid

# **BACTERIOLOGY**

Gram Stains Cultures Sensitivities Strep Screens

Antigen assays (chlamydia, etc.)

H. pylori

# **MYCOBACTERIOLOGY**

Acid Fast Smears Mycobacterial Cultures Sensitivities

# **MYCOLOGY**

**Fungal Cultures** 

DTM

**KOH Preps** 

#### **VIROLOGY**

**RSV** 

HPV assays Cell cultures

# **Endocrinology**

TSH Free T4 Total T4

Trilodothyronine (T3)

T3 Uptake Ferritin Folate PSA B12

Serum-beta-HCG

#### **Toxicology**

Acetaminophen Blood alcohol Carbamazephine

Digoxin
Ethosuximide
Gentamycin
Lithium
Phenobarbitol
Phenytoin
Primidine
Procainamide
NAPA

Quinidine Salicylates Theophylline Tobramycin Valproic acid

# **HEMATOLOGY**

WBC count RBC count Hemoglobin

Hematocrit (Other than spun micro)

Platelet Differential MCV

**Activated Clotting Time** 

Prothrombin time

Partial thromboplastin time

Fibrinogen

Reticulocyte count

Manual WBC by hemocytometer Manual platelet by hemocytometer Manual RBC by hemocytometer

Sperm count

# **IMMUNOHEMATOLOGY**

Rh(D) type Antibody Screening Antibody Identification Compatability testing

ABO group

# **PATHOLOGY**

Dermatopathology Oral pathology PAP smear interpretations Other cytology tests Histopathology

# **RADIOBIOASSAY**

Red cell volume Schilling's test

# **CYTOGENETICS**

Fragile X Buccal smear

#### GUIDELINES FOR COUNTING TESTS FOR CLIA

- o For **histocompatibility**, each HLA typing (including disease associated antigens), HLA antibody screen, or HLA crossmatch is counted as one test.
- o For **microbiology**, susceptibility testing is counted as one test per group of antibiotics used to determine sensitivity for one organism. Cultures are counted as one per specimen regardless of the extent of identification, number of organisms isolated and number of tests/procedures required for identification.
- o Testing for allergens should be counted as one test per individual allergen.
- o For **chemistry** profiles, each individual analyte is counted separately.
- o For **urinalysis**, microscopic and macroscopic examinations, each count as one test. Macroscopics (dipsticks) are counted as one test regardless of the number of reagent pads on the strip.
- o For **complete blood counts**, each <u>measured</u> individual analyte that is ordered <u>and reported</u> is counted separately. Differentials are counted as one test.
- o Do not count calculations (e. g., A/G ratio, MCH, and T7), quality control, quality assurance and proficiency testing assays).
- o For **immunohematology**, each ABO, Rh, antibody screen, crossmatch or antibody identification is counted as one test.
- o For **histopathology**, each block (not slide) is counted as one test. Autopsy services are not included. For those laboratories that perform special stains on histology slides, the test volume is determined by adding the number of special stains performed on slides to the total number of specimen blocks prepared by the laboratory.
- o For **cytology**, each slide (not case) is counted as one test for both Pap smears and nongynecologic cytology.
- o For **cytogenetics**, the number of tests is determined by the number of specimen types processed on each patient; e.g., a bone marrow and a venous blood specimen received on one patient is counted as two tests.